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(54) Solvent extraction of cuprous ions from aqueous solutions

(57) Metal values are recovered from aqueous solutions containing cuprous ion, ferrous ion and halide or pseudo halide ion, especially from solutions derived from leaching of complex sulphide ores with ferric chloride or cupric chloride, by (1) contacting the aqueous solution with a solution in a waterimmiscible organic solvent of a suitable metal extractant, (2) separating the aqueous and organic phases and (3) contacting the organic phase with an aqueous strip solution containing an alkali metal or alkaline earth metal cation and a halide or pseudo-halide anion.

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Process for the extraction of metal values

5 This invention relates to a process for the extraction of metal values from aqueous solutions of metal salts, and in particular to a process for the extraction of copper from aqueous solutions containing cuprous and ferrous cations and halide or pseudo-halide anions.

The use of solvent extraction techniques for the hydrometalurgical recovery of metal values from metal ores has been practised commercially for a number of years. For example copper may be recovered from oxide ores or from ore tailings by treating the crushed ore with sulphuric acid to give an aqueous solution of copper sulphate which is subsequently contacted with a solution in a water-immiscible solvent of a metal extractant whereby the copper values are selectively extracted into the organic solvent phase.

The application of solvent extraction techniques to aqueous solutions containing halide lons has however hitherto presented numerous technical problems.

Of particular importance in this connection is the development of hydrometallurgical routes (as an alternative to smelting) for the extraction of copper from complex sulphide ores (including polymetallic sulphide ores) such as chalcopyrite. It is known to leach such ores using for example aqueous ferric chloride or cupric chloride solution, but the solvent extraction of the resultant leach liquor presents formidable difficulties. Furthermore, whilst there may be process advantages in producing a leach liquor containing copper substantially in the form of cuprous ion, such leach solutions are especially difficult to treat by solvent extraction techniques.

According to the present invention there is provided a process for recovering copper values from an aqueous solution containing cuprous and ferrous cations and halide or pseudo halide anions which comprises the steps of

- (1) contacting the aqueous solution with a solution in a water-immiscible organic solvent of a copper extractant capable of extracting cuprous ion in the presence of halide or pseudo halide ion;
 - (2) separating from the aqueous phase the organic phase containing cuprous ion in the form of a halideor pseudo halide-containing complex with the copper extractant; and
- (3) contacting the organic phase containing the complex of cuprous ion with the extractant with an

 aqueous strip solution containing an alkali metal or alkaline earth metal cation and halide or pseudo-halide

 anion, whereby copper transfers from the organic phase into the aqueous strip solution, the relative

 concentration of halide or pseudo-halide anion being such that the cuprous ion transferred into the aqueous

 strip solution is soluble therein.

The process of the present invention is especially suitable for the treatment of aqueous solutions derived from the leaching of complex sulphide ores using ferric chloride or cupric chloride solutions as leachant. Whilst the leach conditions may be selected to provide a range of ratios of cuprous to cupric ion in the resultant leach liquor, there are certain process advantages in providing a leach liquor containing copper substantially as cuprous ion (for example copper in the form of cuprous ion containing less than 10% of cupric ion). Typical stoichlometry for such a leach processes is represented by the equations:

- (a) $CuFeS_2 + 3FeCl_3 = CuCl + 4FeCl_2 + 2S$
 - (b) $CuFeS_2 + 2CuCl_2 = 4CuCl + FeCl_2 + 2S$
- In the course of the extraction, elemental sulphur is precipitated and is separated from the aqueous leach liquor. The resultant leach liquor, which contains copper substantially in the form of cuprous ion, and Iron substantially in the form of ferrous ion (for example iron in the form of ferrous ion containing less than 10% of ferric ion) may form the feed for the process of the present invention. In general, there will be sufficient ferrous chloride present in solution to prevent precipitation of cuprous chloride, but if necessary additional halide ion may be added to solubilise the cuprous ion. Such halide ion may be added for example in the form of alkali metal or alkaline earth metal chloride. The ferric chloride leachant solution should be sufficiently acid to prevent precipitation of hydrated oxides of Iron, and a typical leach liquor derived from such a process may contain for example 50 gpl cuprous ion, 3M ferrous chloride and 0.2M hydrochloric acid.
- Typical leach conditions using cupric chloride as leachant and typical resultant leach solutions are described for example in United States Patent 3,901,776.

In stage (1) of the process of the present invention, the aqueous solution containing cuprous ion, ferrous ion and halide or pseudo-halide ion, for example a solution derived from the leaching of a complex sulphide ore with ferric chloride solution, is contacted with a solution in a water-immiscible organic solvent of a copper extractant. Suitable copper extractants capable of extracting cuprous ion in the presence of halide or pseudo-halide ion are substituted pyridines described in our European Patent No. 0 057 797. The substituted pyridine extractants described in the European Patent are capable of extracting either cupric or cuprous ions. However, we have found that cuprous ion generally forms a stronger complex with the extractant which is in consequence relatively more difficult to strip. We therefore prefer to use a relatively "weak" extractant, for example an alkyl diester of pyridine-3,5-dicarboxylic acid wherein the alkyl groups of the diester contain a

total of from 16 to 36 carbon atoms, for example the bis isodecyl ester of pyridine 3,5-dicarboxylic acid. Suitable water-immiscible organic solvents are well known in the art, and may be for example aliphatic, aromatic and alicyclic hydrocarbons. The concentration of the extractant in the water-immiscible organic solvent may be chosen to suit the particular leach liquor to be treated. Typical values of extractant 5 concentration in the organic phase are between about 0.1 to 2 Molar, and an especially convenient range is from 0.2 to 0.8 molar in the organic solvent.

The separation step (2) in the process of the present invention may be undertaken using conventional means, for example conventional settlers.

In step (3) in the process of the present invention, the organic solution containing the extractant complex is 10 contacted with an aqueous strip solution containing an alkali metal or alkaline earth metal cation and halide or pseudohalide anion into which the cuprous ion is transferred. The relative concentration of the halide or pseudo-halide anion is such that the cuprous ion is transferred into the aqueous strip solution and is soluble therein. The molarity of halide or pseudo-halide anion required may be readily determined from standard solubility curves for a given concentration of cuprous ion. In the course of the steps (1) to (3), cuprous ion is 15 substantially separated from undesirable impurities originally present in the leach liquor, and from the Iron leachant. The resultant pregnant strip solution contains an increased concentration of purified cuprous ion from which metallic copper may be recovered as described hereafter. Preferably the aqueous strip solution is the barren solution obtained after metallic copper is removed from the pregnant strip solution. In this way one driving force causing the cuprous ion to transfer from the organic extractant phase to the aqueous strip 20 solution is the depletion of copper in the strip solution. An additional driving force may be provided at certain chloride ion concentrations if, as will generally be the case, chloride ion is also removed from pregnant strip solution during the course of the recovery of metallic copper from the cuprous ion.

A variety of methods are available for the recovery of metallic copper from the pregnant strip solution. The details of the recovery method may differ slightly depending on whether a ferric chloride leach or cupric chloride leach is involved. In a first example of such a method, the pregnant strip solution is passed to the cathode compartment of a split electrolytic cell in which the cathode compartment is separated from the anode compartment by an anion exchange membrane. The operation of such a split cell is described in US Department of the Interior Bureau of Mines Report RI 8007. When ferric chloride leachant is used, the aqueous "leach liquor" from which cuprous ion has been removed in the extraction step (1) of the process of 30 the present invention (the raffinate) is passed to the anode compartment. In this way, the cuprous ion is electrodeposited in the cathode compartment as pure cathode copper, whilst the associated halide or pseudo-halide ion transfers through the anion exchange membrane to the anode compartment where the ferrous ion is oxidised to ferric ion. The spent electrolyte from the cathode compartment, which contains a reduced concentration of halide or pseudo-halide ion associated with the removal of cuprous ion, is 35 preferably returned as strip solution to stage (3) of the process of the present invention. Preferably the conditions are selected such that relatively low levels of residual cuprous ion remain in the spent electrolyte since in this way any tendency for cuprous ion to be oxidised to cupric ion may be minimised.

A similar electrolytic process may be used to recover copper and regenerate a cupric chloride leachant. Thus if a split cell is used, the pregnant strip solution may be passed to the cathode compartment, whilst the (impure) cuprous chloride leach liquor may be passed directly to the anode compartment where it is oxidised to cupric solution and the leachant solution is regenerated. Alternatively, the solvent extraction circuit may be arranged so that half of the cuprous ion present in the original leach liquor is retained in the raffinate, the other half being extracted and recovered in the pregnant strip solution. The Pregnant strip solution is then passed to the cathode compartment where copper metal is deposited, whilst the raffinate from the extraction circuit is passed to the anode compartment where it is oxidised to cupric chloride for further use as regenerated leachant. A combination of the process may also be used in which residual cuprous ion in the raffinate is supplemented as appropriate by the original leach liquor before being fed to the anode compartment.

As alternative form of electrolytic cell which may be used to recover metallic copper consists of a 50 conventional cell divided into anode and cathode compartments. The pregnant strip solution is fed to the cell 50 and the cuprous ion is reduced to metallic copper at the cathode, whilst chlorine gas is evolved at the anode, The evolved chlorine gas is preferably used to regenerate the ferrous ion or cuprous ion leachant remaining in the leach liquor after the cuprous ion has been extracted in stage (1) of the process of the present invention (i.e. the raffinate from the extraction stage). The regenerated ferric chloride or cupric chloride leachant may be returned to process further ore. Preferably, the concentration of cuprous ion in the spent 55 electrolyte is relatively low, so that any oxidation of cuprous ion to cupric ion is minimised. A fluidised bed cell may be used if desired to achieve low levels of residual cuprous ion in the spent electrolyte.

In a completely different method of recovering copper from the pregnant strip solution, the strip stage (3) may be operated at elevated temperature, for example about 80°C, and the conditions selected such that the 60. pregnant strip solution is saturated (or supersaturated) with respect to cuprous ion. On cooling the pregnant strip solution to ambient temperature, cuprous chloride precipitates and may be removed by filtration. The filtrate is then returned to the process as barren strip solution. Metallic copper may be recovered from the precipitated cuprous chloride by conventional means, for example fused salt electrolysis or hydrogen reduction. Chlorine evolved during fused salt electrolysis may be used to regenerate ferrous chloride to 65 ferric chloride (or cuprous chloride to cupric chloride) for return to leach further ore. Similarly hydrochloric

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acid evolved during hydrogen reduction may be used in admixture with air to regenerate ferrous ion to ferric.

It will be appreciated that the extraction and strip steps of the process of the present invention may each take place in two or more stages, and that it is preferred to recirculate solutions between the various steps of the process as indicated above to give an integrated overall process. For a truly integrated process additional steps may be incorporated. For example it may be necessary to remove iron introduced into the system from the complex sulphide ore during leaching. Additional oxidation of ferrous iron may also be required to achieve stoichiometry. The removal of excess iron and the oxidation of any residual ferrous ion in the leachant may be achieved for example by air oxidation with the precipitation of hydrated ferric oxide.

10 CLAIMS

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- 1. Process for recovering copper values from an aqueous solution containing cuprous and ferrous cations and halide or pseudo halide anions which comprises the steps of
- (1) contacting the aqueous solution with a solution in a water-immiscible organic solvent of a copper extractant capable of extracting cuprous ion in the presence of halide or pseudo halide ion;
- (2) separating from the aqueous phase the organic phase containing cuprous ion in the form of a halideor pseudo halide-containing complex with the copper extraction; and
- (3) contacting the organic phase containing the complex of cuprous ion with the extractant with an aqueous strip solution containing an alkali metal or alkaline earth metal cation and halide or pseudo-halide anion, whereby copper transfers from the organic phase into the aqueous strip solution, the relative concentration of halide or pseudo-halide anion being such that the cuprous ion transferred into the aqueous strip solution is soluble therein.

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